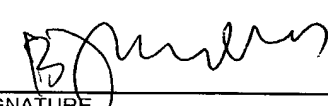


FORM PTO-1390 (REV 11-2000)		U.S. DEPARTMENT OF COMMERCE PATENT AND TRADEMARK OFFICE		ATTORNEY'S DOCKET NUMBER <b>3557-13</b>	
<b>TRANSMITTAL LETTER TO THE UNITED STATES DESIGNATED/ELECTED OFFICE (DO/EO/US) CONCERNING A FILING UNDER 35 U.S.C. 371</b>				U.S. APPLICATION NO. (If known, see 37 C.F.R. 1.5) <b>10/089370</b> Unknown	
INTERNATIONAL APPLICATION NO. <b>PCT/EP00/09839</b>		INTERNATIONAL FILING DATE <b>7 October 2000</b>		PRIORITY DATE CLAIMED <b>11 October 1999</b>	
TITLE OF INVENTION <b>PLANT PRPP AMIDOTRANSFERASE</b>					
APPLICANT(S) FOR DO/EO/US <b>LERCHL et al</b>					
Applicant herewith submits to the United States Designated/Elected Office (DO/EO/US) the following items and other information:					
1. <input checked="" type="checkbox"/> This is a <b>FIRST</b> submission of items concerning a filing under 35 U.S.C. 371. 2. <input type="checkbox"/> This is a <b>SECOND</b> or <b>SUBSEQUENT</b> submission of items concerning a filing under 35 U.S.C. 371. 3. <input checked="" type="checkbox"/> This is an express request to begin national examination procedures (35 U.S.C. 371(f)). The submission must include items (5), (6), (9) and (21) indicated below. 4. <input checked="" type="checkbox"/> The U.S. has been elected by the expiration of 19 months from the priority date (Article 31). 5. A copy of the International Application as filed (35 U.S.C. 371(c)(2)). a. <input type="checkbox"/> is attached hereto (required only if not communicated by the International Bureau). b. <input checked="" type="checkbox"/> has been communicated by the International Bureau. c. <input type="checkbox"/> is not required, as the application was filed in the United States Receiving Office (RO/US). 6. <input checked="" type="checkbox"/> An English language translation of the International Application as filed (35 U.S.C. 371(c)(2)). a. <input checked="" type="checkbox"/> is attached hereto. b. <input type="checkbox"/> has been previously submitted under 35 U.S.C. 154(d)(4). 7. <input type="checkbox"/> Amendments to the claims of the International Application under PCT Article 19 (35 U.S.C. 371(c)(3)). a. <input type="checkbox"/> are attached hereto (required only if not communicated by the International Bureau). b. <input type="checkbox"/> have been communicated by the International Bureau. c. <input type="checkbox"/> have not been made; however, the time limit for making such amendments has <b>NOT</b> expired. d. <input type="checkbox"/> have not been made and will not be made. 8. <input checked="" type="checkbox"/> An English language translation of the amendments to the claims under PCT Article 19 (35 U.S.C. 371(c)(3)). 9. <input checked="" type="checkbox"/> An oath or declaration of the inventor(s) (35 U.S.C. 371(c)(4)). 10. <input checked="" type="checkbox"/> A English language translation of the annexes of the International Preliminary Examination Report under PCT Article 36 (35 U.S.C. 371(c)(5)).					
<b>Items 11 To 20 below concern document(s) or information included:</b>					
11. <input checked="" type="checkbox"/> An Information Disclosure Statement under 37 C.F.R. 1.97 and 1.98. 12. <input checked="" type="checkbox"/> An assignment document for recording. A separate cover sheet in compliance with 37 C.F.R. 3.28 and 3.31 is included. 13. <input checked="" type="checkbox"/> A FIRST preliminary amendment. 14. <input type="checkbox"/> A SECOND or SUBSEQUENT preliminary amendment. 15. <input type="checkbox"/> A substitute specification. 16. <input type="checkbox"/> A change of power of attorney and/or address letter. 17. <input checked="" type="checkbox"/> A computer-readable form of the sequence listing in accordance with PCT Rule 13ter.2 and 35 U.S.C. 1.821-1.825. 18. <input type="checkbox"/> A second copy of the published international application under 35 U.S.C. 154(d)(4). 19. <input type="checkbox"/> A second copy of the English language translation of the international application under 35 U.S.C. 154(d)(4). 20. <input checked="" type="checkbox"/> Other items or information. Paper copy of sequence listing					

U.S. APPLICATION NO. (If known, see 37 C.F.R. 1.492(e)) <b>10/089370</b> Unknown		INTERNATIONAL APPLICATION NO. <b>PCT/EP00/09839</b>		ATTORNEY'S DOCKET NUMBER <b>3557-13</b>	
21. <input checked="" type="checkbox"/> The following fees are submitted:				<b>CALCULATIONS</b> PTO USE ONLY	
<b>BASIC NATIONAL FEE (37 C.F.R. 1.492(a)(1)-(5)):</b> -- Neither international preliminary examination fee (37 C.F.R. 1.482) nor international search fee (37 C.F.R. 1.445(a)(2)) paid to USPTO and International Search Report not prepared by the EPO or JPO .....\$1040.00 -- International preliminary examination fee (37 C.F.R. 1.482) not paid to USPTO but International Search Report prepared by the EPO or JPO .....\$890.00 -- International preliminary examination fee (37 C.F.R. 1.482) not paid to USPTO but international search fee (37 C.F.R. 1.445(a)(2)) paid to USPTO .....\$740.00 -- International preliminary examination fee (37 C.F.R. 1.482) paid to USPTO but all claims did not satisfy provisions of PCT Article 33(1)-(4) .....\$710.00 -- International preliminary examination fee (37 C.F.R. 1.482) paid to USPTO and all claims satisfied provisions of PCT Article 33(1)-(4) .....\$100.00					
ENTER APPROPRIATE BASIC FEE AMOUNT =					
Surcharge of \$130.00 for furnishing the oath or declaration later than <input type="checkbox"/> 20 <input type="checkbox"/> 30 months from the earliest claimed priority date (37 C.F.R. 1.492(e)).					
				\$	890.00
				\$	0.00
CLAIMS	NUMBER FILED	NUMBER EXTRA	RATE		
Total Claims	17	-20 =	0	X	\$18.00
Independent Claims	5	-3 =	2	X	\$84.00
MULTIPLE DEPENDENT CLAIMS(S) (if applicable)			\$280.00	\$	0.00
<b>TOTAL OF ABOVE CALCULATIONS =</b>				\$	<b>1058.00</b>
<input type="checkbox"/> Applicant claims small entity status. See 37 CFR 1.27. The fees indicated above are reduced by 1/2.				\$	0.00
<b>SUBTOTAL =</b>				\$	<b>1058.00</b>
Processing fee of \$130.00, for furnishing the English Translation later than <input type="checkbox"/> 20 <input type="checkbox"/> 30 months from the earliest claimed priority date (37 C.F.R. 1.492(f)).				\$	0.00
<b>TOTAL NATIONAL FEE =</b>				\$	<b>1058.00</b>
Fee for recording the enclosed assignment (37 C.F.R. 1.21(h)). The assignment must be accompanied by an appropriate cover sheet (37 C.F.R. 3.28, 3.31). \$40.00 per property				\$	40.00
Fee for Petition to Revive Unintentionally Abandoned Application (\$1280.00 - Small Entity = \$640.00)				\$	0.00
<b>TOTAL FEES ENCLOSED =</b>				\$	<b>1098.00</b>
				Amount to be:	
				refunded	\$
				Charged	\$
a. <input checked="" type="checkbox"/> A check in the amount of \$1098.00 to cover the above fees is enclosed. b. <input type="checkbox"/> Please charge my Deposit Account No. 14-1140 in the amount of \$_____ to cover the above fees. A duplicate copy of this form is enclosed. c. <input checked="" type="checkbox"/> The Commissioner is hereby authorized to charge any additional fees which may be required, or credit any overpayment to Deposit Account No. 14-1140. A duplicate copy of this form is enclosed. d. <input checked="" type="checkbox"/> The entire content of the foreign application(s), referred to in this application is/are hereby incorporated by reference in this application.					
<b>NOTE:</b> Where an appropriate time limit under 37 C.F.R. 1.494 or 1.495 has not been met, a petition to revive (37 C.F.R. 1.137(a) or (b)) must be filed and granted to restore the application to pending status.					
<b>SEND ALL CORRESPONDENCE TO:</b> NIXON & VANDERHYE P.C. 1100 North Glebe Road, 8 <sup>th</sup> Floor Arlington, Virginia 22201-4714 Telephone: (703) 816-4000					
				 SIGNATURE	
				<b>B. J. Sadoff</b> NAME	
				<b>36,663</b> REGISTRATION NUMBER	
				<b>March 29, 2002</b> Date	

**IN THE UNITED STATES PATENT AND TRADEMARK OFFICE**

In re Patent Application of

**Lerchl et al.**

Atty. Ref.: **3557-13**

Serial No. **Unassigned**

Group: **Unassigned**

National Phase of: **PCT/EP00/09839**

International Filing Date: **7 October 2000**

Filed: **March 29, 2002**

Examiner: **Unassigned**

For: **PLANT PRPP AMIDOTRANSFERASE**

\* \* \* \* \*

**March 29, 2002**

Assistant Commissioner for Patents  
Washington, DC 20231

Sir:

**PRELIMINARY AMENDMENT**

Prior to calculation of the filing fee and in order to place the above identified application in better condition for examination, please amend as follows:

**IN THE SPECIFICATION**

Page 1, after the title insert the following:

-- This application is the US national phase of international application PCT/EP00/09839 filed 7 October 2000, which designated the U.S. --.

Insert the attached Sequence Listing in place of the originally-filed Sequence Listing.

**IN THE CLAIMS**

Please substitute the following amended claims for corresponding claims previously presented. A copy of the amended claims showing current revisions is attached.



Lerchl et al.  
Serial No. **Unassigned**

**REMARKS**

Attached hereto is a marked-up version of the changes made to the claims by the current amendment. The attached page is captioned "**Version with markings to show changes made.**"

The attached paper and computer readable copies of the Sequence Listing are the same. No new matter has been added. A separate Statement to this effect is attached.

The above amendments are made to place the claims in a more traditional format.

Respectfully submitted,

**NIXON & VANDERHYE P.C.**

By:



**B. J. Sadoff**

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**Lerchl et al.**  
Serial No. **Unassigned**

**VERSION WITH MARKINGS TO SHOW CHANGES MADE**

6. (Amended) The use of a DNA sequence as claimed in claim 1 [or 2] for introduction into pro- or eukaryotic cells, this sequence optionally being linked to control elements which ensure transcription and translation in the cells and leading to the expression of a translatable mRNA which causes the synthesis of a plant PRPP amidotransferase.

7. (Amended) The use of a DNA sequence as claimed in claim 1 [or 2] for generating an assay system for identifying herbicidally active plant PRPP amidotransferase inhibitors.

8. (Amended) A method of finding herbicidally active substances which inhibit the activity of the plant PRPP amidotransferase, which comprises preparing, in a first step, PRPP amidotransferase using a DNA sequence as claimed in claim 1 [or 2] and measuring, in a second step, the activity of the plant PRPP amidotransferase in the presence of a test substance.

11. (Amended) An assay system based on the expression of a DNA sequence SEQ-ID No. 1 or SEQ-ID No.9 as claimed in claim 1 [or 2] for identifying herbicidally active plant PRPP amidotransferase inhibitors.

14. (Amended) A plant PRPP amidotransferase inhibitor identified using an assay system as claimed in claim 11 [or 12].

15. (Amended) An inhibitor as claimed in claim 13 [or 14] for use as herbicide.

Plant PRPP amidotransferase

The present invention relates to the identification of plant PRPP  
5 amidotransferase (phosphoribosyl-pyrophosphate amidotransferase,  
E.C. 2.4.2.14) as novel target for herbicidal active ingredients.  
The present invention furthermore relates to DNA sequences  
encoding a polypeptide with PRPP amidotransferase activity.  
Moreover, the invention relates to the use of a nucleic acid  
10 encoding a protein with PRPP amidotransferase activity which  
originates from plants for generating an assay system for  
identifying herbicidally active PRPP amidotransferase inhibitors  
and to plant PRPP amidotransferase identified using this assay  
system. The invention furthermore relates to the use of the  
15 nucleic acid SEQ-ID No. 1 or SEQ-ID No. 3 encoding plant PRPP  
amidotransferase for the generation of plants with an increased  
resistance to PRPP amidotransferase inhibitors and for the  
generation of plants with a modified purine nucleotide content.  
Moreover, the invention relates to a method of eliminating  
20 undesired vegetation, where the plants to be eliminated are  
treated with a compound which binds specifically to PRPP  
amidotransferase encoded by a DNA sequence SEQ-ID No 1 or a DNA  
sequence which hybridizes with this DNA sequence, and inhibits  
its function.

25

Plants are capable of synthesizing their cell components from  
carbon dioxide, water and inorganic salts.

This process is only possible by exploiting biochemical reactions  
30 for synthesizing organic substances. Nucleotides are synthesized  
de novo in plants. Being components of the nucleic acids, they  
are particularly important. Covalently bound, nucleotides  
activate carbohydrates for polysaccharide biosynthesis. They  
furthermore activate head groups for lipid biosynthesis.  
35 Nucleotides are involved in virtually all metabolic pathways.  
Nucleoside triphosphates, especially ATP, drive most of the  
energy-requiring reactions of the cell. Adenine nucleotides are  
additionally also found as components in essential factors such  
as coenzyme A and in nicotinamide and flavin coenzymes, which are  
40 involved in a large number of cellular reactions. The coupled  
hydrolysis of guanosine-5'-triphosphate (GTP) defines a direction  
of reaction for various cellular processes such as protein  
translation, assembly of microtubuli, vesicular transport, signal  
transduction and cell division. Furthermore, nucleotides  
45 constitute the starting metabolites for the biosynthesis of

methylxanthines such as caffeine and theobromine in the plant family of the Rubiaceae and Theaceae.

cDNAs which encode PRPP amidotransferase have been isolated and characterized from various bacterial, animal and vegetable organisms. Plant PRPP amidotransferase cDNAs have been isolated via complementation of *E. coli* purF mutants and via DNA hybridization techniques from *Glycine max*, *Vigna aconitifolia* and from *Arabidopsis thaliana* (Ito et al., Plant Molecular Biology 26(1994), 529-533; Kim et al., The Plant Journal 7(1995), 77-86). Sequence homology suggests that the encoded enzymes as well as the *E. coli* PRPP amidotransferase contain 4Fe-4S clusters. The plant PRPP amidotransferase amino acid sequences, which in comparison with *E. coli* are extended at the N terminus, show similarity to plastid signal sequences.

20 Several PRPP amidotransferase isoenzymes which are expressed differentially are found in plants. The RNA for *Arabidopsis thaliana* AtATase1, for example, accumulates preferentially in the roots, while the AtATase2 transcripts are found predominantly in young leaves and flowers (Ito et al., Plant Molecular Biology  
25 26(1994), 529-533). In *Vigna aconitifolia*, a PRPP amidotransferase RNA accumulates mainly in root nodules and is induced in root tissues by L-Glutamine (Kim et al., The Plant Journal 7(1995), 77-86).

30 Since plants depend on an effective nucleotide metabolism, it can be assumed that the enzymes which are involved in nucleotide biosynthesis are suitable as target for herbicides. Thus, there have already been described active ingredients which inhibit *de novo* purine biosynthesis in plants. An example which may be  
35 mentioned is the natural substance hydanthocidin, which, after phosphorylation *in planta* inhibits adenylosuccinate synthetase (ASS); (Siehl et al., Plant Physiol. 110(1996), 753-758).

Inhibitors for enzymes of purine biosynthesis are, moreover, also known for their pharmacological action in animals and microorganisms: folate analogs inhibit, inter alia, the enzyme GAR transformylase and have an antiproliferative, antiinflammatory and immunosuppressant action. Mycophenolic acid (MPA), an IMP dehydrogenase inhibitor in the GMP synthetic pathway, has an antimicrobial, antiviral and immunosuppressant action (Kitchin et al, Journal of the American Academy of



0050/50796

## 3

Dermatology 37(1997), 445-449).

Bacterial PRPP amidotransferase can be inhibited for example by glutamine antagonists such as, for example, azaserine,  
5 6-diazo-5-oxo-L-norleucine (DON) or L-2-amino-4-oxo-5-chloropentanoic acid and by mercaptopurine and thioguanosine. Glutamine antagonists are not specific to PRPP amidotransferase and also affect other purine biosynthesis enzymes, such as formylglycinamidine ribotide synthase. The  
10 efficacy of glutamine antagonists on plant PRPP amidotransferase is still to be proven.

It is an object of the present invention to provide proof that PRPP amidotransferase in plants is a suitable herbicidal target,  
15 to isolate a complete plant cDNA encoding the enzyme PRPP amidotransferase and functionally express it in bacterial or eukaryotic cells, and to produce an efficient and simple PRPP amidotransferase assay system for carrying out inhibitor-enzyme binding studies.

20 We have found that this object is achieved by the isolation of genes which encode the plant enzyme PRPP amidotransferase, the generation of PRPP amidotransferase antisense constructs, and the functional expression of PRPP amidotransferase in bacterial or  
25 eukaryotic cells.

It is an object of the present invention to isolate full-length cDNAs encoding functional PRPP amidotransferase (E.C.2.4.2.14) from tobacco (*Nicotiana tabacum*).

30 A first subject-matter of the present invention is a DNA sequence SEQ-ID NO. 1 or SEQ-ID NO. 3 containing the encoding region of a plant PRPP amidotransferase from tobacco, see Example 1.

35 Another subject-matter of the invention is DNA sequences which are derived from SEQ-ID NO. 1 or SEQ-ID NO. 3 or which hybridize with one of these sequences and which encode a protein which has the biological activity of a PRPP amidotransferase.

40 Tobacco plants of the line *Nicotiana tabacum* cv. Samsun NN which carry a PRPP amidotransferase antisense construct have been characterized in greater detail. The plants show different degrees of retarded growth and bleaching of the leaves. The transgenic lines and the progeny of the 1<sup>st</sup> and 2<sup>nd</sup> generation  
45 showed a reduced growth in soil. Using Northern hybridization, it was detected that the RNA quantity of PRPP amidotransferase was reduced in plants with reduced growth compared with the wild

0050/50796

4

type. Furthermore, measurement of the enzyme activity detected that the amount of PRPP amidotransferase activity was reduced in the transgenic lines compared with wild-type plants, see Example 7. Growth retardation and the reduction in PRPP amidotransferase activity correlate. This clear connection identifies PRPP amidotransferase for the first time unambiguously as suitable target protein for herbicidal active ingredients.

To be able to find efficient inhibitors of plant PRPP amidotransferase, it is necessary to provide suitable assay systems with which inhibitor/enzyme binding studies can be carried out. To this end, for example, the complete cDNA sequence of tobacco PRPP amidotransferase is cloned into an expression vector (pQE, Qiagen) and overexpressed in E. coli, see Example 2.

Alternatively, however, it is possible to express the expression cassette containing a DNA sequence of SEQ-ID No. 1 or SEQ-ID NO. 3 for example in other bacteria, in yeasts, fungi, algae, plant cells, insect cells or mammalian cells, see Example 4.

The PRPP amidotransferase protein which is expressed with the aid of the expression cassette according to the invention is particularly suitable for finding inhibitors which are specific to PRPP amidotransferase.

To this end, for example, the plant PRPP amidotransferase can be employed in an enzyme assay in which the PRPP amidotransferase activity is determined in the presence and absence of the active ingredient to be tested. A comparison of the two activity determinations allows a qualitative and quantitative statement to be made on the inhibitory behavior of the active ingredient to be tested, see Example 3.

The assay system according to the invention allows a multiplicity of chemicals to be tested rapidly and simply for herbicidal properties. Using this method, substances with a potent action can be selected specifically and reproducibly from amongst a large number of substances, in order that further in-depth tests with which the skilled worker is familiar are carried out subsequently with these substances.

The invention furthermore relates to a method of identifying herbicidally active substances which inhibit the PRPP amidotransferase activity in plants, with the following steps:

45

5

- a) the generation of transgenic plants, plant tissues or plant cells which comprise an additional DNA sequence encoding an enzyme with PRPP amidotransferase activity and which are capable of overexpressing an enzymatically active PRPP amidotransferase;
- b) applying a substance to transgenic plants, plant cells, plant tissue or plant parts and to untransformed plants, plant cells, plant tissue or plant parts;
- c) determining the growth or the viability of the transgenic and the untransformed plants, plant cells, plant tissue or plant parts after application of the chemical substance; and
- d) comparing the growth or the viability of the transgenic and the untransformed plants, plant cells, plant tissue or plant parts after applying the chemical substance;

where a suppression of the growth or the viability of the  
20 untransformed plants, plant cells, plant tissue or plant parts,  
but an absence of potent suppression of the growth or viability  
of the transgenic plants, plant cells, plant tissue or plant  
parts, confirms that the substance of b) shows herbicidal  
activity and inhibits the PRPP amidotransferase enzyme activity  
25 in plants.

Another subject-matter of the invention is a method of identifying plant PRPP amidotransferase inhibitors with a potentially herbicidal action by cloning the gene of a plant PRPP amidotransferase, overexpressing it in a suitable expression cassette - for example in insect cells - disrupting the cells and employing the cell extract in an assay system for measuring the enzyme activity in the presence of low-molecular-weight chemicals, either directly or after concentration or isolation of the enzyme PRPP amidotransferase.

Another subject-matter of the invention is compounds with a herbicidal action which can be identified with the above-described assay system.

The invention furthermore relates to a method of eliminating undesired vegetation, where the plants to be eliminated are treated with a compound which binds specifically to plant PRPP amidotransferase and inhibits its function.

## 6

Herbicidally active PRPP amidotransferase inhibitors can be employed as defoliants, desiccants, haulm killers and, in particular, as herbicides. Weeds in the widest sense are to be understood as meaning all plants which grow in locations where  
5 they are undesired. Whether the active ingredients found with the aid of the assay system according to the invention act as total or selective herbicides depends, inter alia, on the quantity applied.

10 Herbicidally active PRPP amidotransferase inhibitors can be used, for example, against the following weeds:

Dicotyledonous weeds of the genera:

Sinapis, Lepidium, Galium, Stellaria, Matricaria, Anthemis,  
15 Galinsoga, Chenopodium, Urtica, Senecio, Amaranthus, Portulaca, Xanthium, Convolvulus, Ipomoea, Polygonum, Sesbania, Ambrosia, Cirsium, Carduus, Sonchus, Solanum, Rorippa, Rotala, Lindernia, Lamium, Veronica, Abutilon, Emex, Datura, Viola, Galeopsis, Papaver, Centaurea, Trifolium, Ranunculus, Taraxacum.

20

Monocotyledonous weeds of the genera:

Echinochloa, Setaria, Panicum, Digitaria, Phleum, Poa, Festuca, Eleusine, Brachiaria, Lolium, Bromus, Avena, Cyperus, Sorghum, Agropyron, Cynodon, Monochoria, Fimbristylis, Sagittaria,  
25 Eleocharis, Scirpus, Paspalum, Ischaemum, Sphenoclea, Dactyloctenium, Agrostis, Alopecurus, Apera.

Subject-matter of the invention are also expression cassettes whose sequence encodes a tobacco PRPP amidotransferase or its  
30 functional equivalent. The nucleic acid sequence can be, for example, a DNA or a cDNA sequence.

In addition, the expression cassettes according to the invention comprise regulatory nucleic acid sequences which govern the  
35 expression of the encoding sequence in the host cell. In accordance with a preferred embodiment, an expression cassette according to the invention encompasses upstream, i.e. at the 5' end of the encoding sequence, a promoter, and downstream, i.e. at the 3' end, a polyadenylation signal and, if appropriate, other  
40 regulatory elements which are operatively linked to the encoding sequence for the PRPP amidotransferase gene, which sequence lies between the promoter and the polyadenylation signal. Operative linkage is to be understood as meaning the sequential arrangement of promoter, encoding sequence, terminator and, if appropriate,  
45 other regulatory elements in such a manner that each of the

0050/50796

7

regulatory elements can function as intended when the encoding sequence is expressed.

An expression cassette according to the invention is generated by  
 5 fusing a suitable promoter with a suitable PRPP amidotransferase  
 DNA sequence and a polyadenylation signal using customary  
 recombination and cloning techniques as they are described, for  
 example, by T. Maniatis, E.F. Fritsch and J. Sambrook, Molecular  
 Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory, Cold  
 10 Spring Harbor, NY (1989) and by T.J. Silhavy, M.L. Berman and  
 L.W. Enquist, Experiments with Gene Fusions, Cold Spring Harbor  
 Laboratory, Cold Spring Harbor, NY (1984) and by Ausubel, F.M. et  
 al., Current Protocols in Molecular Biology, Greene Publishing  
 Assoc. and Wiley-Interscience (1987).

15 Subject-matter of the invention are also functionally equivalent  
 DNA sequences which encode a PRPP amidotransferase gene and which  
 show a sequence homology with the DNA sequence SEQ-ID No. 1 or  
 SEQ-ID No. 3 of 40 to 100%, based on the total length of the DNA  
 20 sequence.

Preferred subject-matter of the invention are functionally  
 equivalent DNA sequences which encode a PRPP amidotransferase  
 gene and which show a sequence homology with the DNA sequence  
 25 SEQ-ID No. 1 or SEQ-ID No. 3 of 60 to 100%, based on the total  
 length of the DNA sequence.

Particularly preferred subject-matter of the invention are  
 functionally equivalent DNA sequences which encode a PRPP  
 30 amidotransferase gene and which show a sequence homology with the  
 DNA sequence SEQ-ID No. 1 or SEQ-ID No. 3 of 80 to 100%, based on  
 the total length of the DNA sequence.

Functionally equivalent sequences which encode a PRPP  
 35 amidotransferase gene are in accordance with the invention those  
 sequences which retain the desired functions, despite a deviating  
 nucleotide sequence. Functional equivalents thus encompass  
 naturally occurring variants of the sequences described herein,  
 but also artificial nucleotide sequences, for example those which  
 40 have been obtained by chemical synthesis and which are adapted to  
 suit the codon usage of a plant.

A functional equivalent is also to be understood as meaning in  
 particular natural or artificial mutations of an originally  
 45 isolated sequence which encodes a PRPP amidotransferase and which  
 continues to show the desired function. Mutations encompass  
 substitutions, additions, deletions, exchanges or insertions of

## 8

one or more nucleotide residues. Thus, the present invention for example also extends to those nucleotide sequences which are obtained by modifying this nucleotide sequence. The target of such a modification can be, for example, the further delimitation of the encoding sequence contained therein or else, for example, the introduction of further restriction enzyme cleavage sites.

Functionelle equivalents are also those variants whose function is reduced or increased compared with the starting gene or gene fragment.

In addition, the expression cassette according to the invention can also be employed for the transformation of bacteria, cyanobacteria, yeasts, filamentous fungi and algae, with the purpose of producing sufficient amounts of the enzyme PRPP amidotransferase.

Another subject-matter of the invention is a tobacco protein characterized by the amino acid sequence SEQ-ID NO: 2 or SEQ-ID NO. 4 or derivatives or parts of this protein with PRPP amidotransferase activity.

Subject-matter of the invention are also plant proteins with PRPP amidotransferase activity with an amino acid sequence homology to the tobacco PRPP amidotransferase with the SEQ-ID NO: 2 or SEQ-ID NO. 4 of 20 - 100% identity.

Preferred are plant proteins with PRPP amidotransferase activity with an amino acid sequence homology to the tobacco PRPP amidotransferase with the sequences SEQ-ID NO: 2 or SEQ-ID NO. 4 of 50 - 100% identity.

Particularly preferred are plant proteins with PRPP amidotransferase activity with an amino acid sequence homology to the tobacco PRPP amidotransferases with the sequences SEQ-ID NO: 2 or SEQ-ID NO. 4 of 80 - 100% identity.

It was another object of the invention to overexpress the PRPP amidotransferase gene in plants in order to generate plants which tolerate PRPP amidotransferase inhibitors.

Overexpression, in a plant, of the gene sequence SEQ-ID NO. 1 or SEQ-ID NO. 3, which encodes a PRPP amidotransferase, results in an increased resistance to PRPP amidotransferase inhibitors. The transgenic plants generated thus are also subject-matter of the invention.

0050/50796

## 9

Expressional efficacy of the recombinantly expressed PRPP amidotransferase gene can be determined, for example, in vitro by shoot-meristem propagation or by a germination test. Moreover, the expression of a PRPP amidotransferase gene which has been  
5 altered in terms of type and level, and its effects on the resistance to PRPP amidotransferase inhibitors can be tested in greenhouse experiments using test plants.

Subject-matter of the invention are also transgenic plants,  
10 transformed with an expression cassette according to the invention containing the DNA sequence SEQ-ID No. 1 or SEQ-ID No. 3, which have been made tolerant to PRPP amidotransferase inhibitors by additionally expressing the DNA sequence SEQ-ID No. 1 or SEQ-ID No. 3, and transgenic cells, tissues, parts and  
15 propagation material of such plants. Especially preferred in this context are transgenic crop plants such as, for example, barley, wheat, rye, maize, soya, rice, cotton, sugar beet, canola, sunflowers, flax, hemp, potatoes, tobacco, tomatoes, oilseed rape, alfalfa, lettuce and the various tree, nut and grapevine  
20 species, and also legumes.

A change in the nucleotide content in plants may be useful under various circumstances. For example, nucleotides are added to plant-based baby formulas to achieve a nutrient composition which  
25 corresponds to breast milk. Furthermore, an optimized nucleotide content would be helpful when patients are fed by gastric tube. A reduced purine nucleotide content in nutritional plants is relevant for the dietetic diet of patients suffering from gout. Furthermore, nucleotides make and enhance flavors, so that an  
30 altered nucleotide content has an effect on the palatability of plants.

Another subject-matter of the invention are thus plants which, following expression of the DNA sequence SEQ-ID No. 1 or SEQ-ID  
35 No. 3 in the plant, have a modified purine nucleotide content. It is preferred to increase the content of the purine nucleotides IMP, AMP and/or GMP, or of their di- or trinucleotides ADP, ATP or GDP and GTP.

40 A plant with a modified purine nucleotide content is generated, for example, by expressing, in the plant, an additional DNA sequence SEQ-ID No. 1 or SEQ-ID No. 3 in sense or antisense orientation. A modified purine nucleotide content means that both plants with an increased purine nucleotide content (in the case  
45 of sense orientation) and plants with a reduced guanosine nucleotide content (in the case of sense orientation

## 10

[cosuppression] or antisense orientation) can be generated.

An increased purine nucleotide content means for the purposes of the present invention for example the artificially acquired  
5 ability of an increased purine nucleotide biosynthesis rate by functionally overexpressing the PRPP amidotransferase gene in the plant in comparison with the non-recombinant plant for the duration of at least one plant generation.

10 Another subject-matter of the invention is the use of plant PRPP amidotransferase for altering the methylxanthine concentration in plants.

Particularly preferred are sequences which ensure targeting into  
15 the apoplast, into plastids, into the vacuole, into the mitochondrion, into the endoplasmatic reticulum (ER), or which, owing to the absence of suitable operative sequences, ensure that the product remains in the compartment where it is formed, in the cytosol, (Kermode, Crit. Rev. Plant Sci. 15, 4 (1996), 285-423).

20 For example, the plant expression cassette can be introduced into the plant transformation vector pBinAR, see Example 5.

A suitable promoter of the expression cassette according to the  
25 invention is, in principle, any promoter which is capable of governing the expression of foreign genes in plants. It is preferred to use, in particular, a plant promoter or a promoter derived from a plant virus. Particularly preferred is the cauliflower mosaic virus CaMV 35S promoter (Franck et al., Cell  
30 21(1980), 285-294). This promoter contains different recognition sequences for transcriptional effectors which, in their totality, lead to permanent and constitutive expression of the gene which has been introduced (Benfey et al., EMBO J., 8 (1989), 2195-2202).

35 The expression cassette according to the invention may also comprise a chemically inducible promoter which allows expression of the exogenous PRPP amidotransferase gene in the plant to be governed at a particular point in time. Such promoters which are  
40 described in the literature and which can be used are, inter alia, for example the PRP1 promoter (Ward et al., Plant. Mol. Biol. (1993) 22, 361-366), a salicylic acid-inducible promoter (WO 95/19443), a benzenesulfonamide-inducible promoter (EP 388186), a tetracyclin-inducible promoter (Gatz et al., Plant J.  
45 (1992) 2, 397-404), an abscisic acid-inducible promoter (EP0335528) or an ethanol- or cyclohexanone-inducible promoter



0050/50796

11

(WO 93/21334).

Particularly preferred promoters are furthermore those which ensure expression in tissues or parts of the plant in which the biosynthesis of purines or their precursors takes place. Promoters which ensure leaf-specific expression must be mentioned in particular. Promoters which must be mentioned are the potato cytosolic FBPase or the potato ST-LSI promoter (Stockhaus et al., EMBO J., 8 (1989) 2445-245).

10

A foreign protein can be expressed stably in the seeds of transgenic tobacco plants to an extent of 0.67% of the total soluble seed protein with the aid of a seed-specific promoter (Fiedler and Conrad, Bio/Technology 10 (1995), 1090-1094). The expression cassette according to the invention can therefore contain, for example, a seed-specific promoter (preferably the phaseolin promoter, the USP promoter or the LEB4 promoter), the LEB4 signal peptide, the gene to be expressed and an ER retention signal.

20

The inserted nucleotide sequence encoding a PRPP amidotransferase can be produced synthetically or obtained naturally or contain a mixture of synthetic and natural DNA components. In general, synthetic nucleotide sequences are generated with codons which are preferred by plants. These codons which are preferred by plants can be determined from codons with the highest protein frequency expressed in the plant species of the highest interest. When preparing an expression cassette, a variety of DNA fragments may be manipulated in order to obtain a nucleotide sequence which expediently reads in the correct direction and which is equipped with a correct reading frame. Adaptors or linkers can be added to the fragments in order to link the DNA fragments to each other.

Other suitable DNA sequences are artificial DNA sequences as long as they mediate the desired property by increasing the purine nucleotide content in the plant by overexpressing the PRPP amidotransferase gene in crop plants, as described above by way of example. Such artificial DNA sequences can be determined for example by backtranslating of proteins which have PRPP amidotransferase activity and which have been constructed by means of molecular modeling, or they can be determined by in vitro selection. Especially suitable are encoding DNA sequences which have been obtained by backtranslating a polypeptide sequence in accordance with the host-plant-specific codon usage. The specific codon usage can be determined readily by a skilled worker familiar with methods of plant genetics by means of

Other suitable equivalent nucleic acid sequences according to the  
5 invention which must be mentioned are sequences which encode  
fusion proteins, the component of the fusion protein being a  
plant PRPP amidotransferase polypeptide or a functionally  
equivalent part thereof. The second part of the fusion protein  
can be, for example, another polypeptide with enzymatic activity  
10 or an antigenic polypeptide sequence, with the aid of which  
detection of PRPP amidotransferase expression is possible (for  
example myc-tag or his-tag). However, it is preferably a  
regulatory protein sequence such as, for example, a signal or  
transit peptide, which leads the PRPP amidotransferase protein to  
15 the desired site of action.

The promoter and terminator regions according to the invention should expediently be provided, in the direction of transcription, with a linker or polylinker containing one or more  
20 restriction sites for insertion of this sequence. As a rule, the linker has 1 to 10, in most cases 1 to 8, preferably 2 to 6, restriction sites. In general, the linker within the regulatory regions has a size less than 100 bp, frequently less than 60 bp, but at least 5 bp. The promoter according to the invention may be  
25 native, or homologous, or else foreign, or heterologous, to the host plant. The expression cassette according to the invention comprises, in the 5'-3' direction of transcription, the promoter according to the invention, any sequence and a region for transcriptional termination. Various termination regions can be  
30 exchanged for each other as desired.

Manipulations which provide suitable restriction cleavage sites or which eliminate the excess DNA or restriction cleavage sites may also be employed. *In vitro* mutagenesis, primer repair, 35 restriction or ligation may be used in cases where insertions, deletions or substitutions such as, for example, transitions and transversions, are suitable. Complementary ends of the fragments may be provided for ligation in the case of suitable manipulations such as, for example, restriction, chewing-back or 40 filling overhangs for blunt ends.

Preferred polyadenylation signals are plant polyadenylation signals, preferably those which correspond essentially to *Agrobacterium tumefaciens* T-DNA polyadenylation signals, in particular those of the gene 3' of the T-DNA (octopine synthase) of the Ti plasmid pTiACH5 (Gielen et al., EMBO J., 3 (1984),



0050/50796

## 14

In addition, constitutive expression of the exogenous PRPP amidotransferase gene is advantageous. On the other hand, inducible expression may also be desirable.

- 5 Using the recombination and cloning techniques cited above, the expression cassettes according to the invention can be cloned into suitable vectors which allow them to be multiplied, for example in *E. coli*. Suitable cloning vectors are, inter alia, pBR332, pUC series, M13mp series and pACYC184. Especially
- 10 suitable are binary vectors which are capable of replication both in *E. coli* and in agrobacteria.

- Another subject-matter of the invention relates to the use of an expression cassette according to the invention for transforming
- 15 plants, plant cells, plant tissues or parts of plants. The preferred purpose of the use is to increase the PRPP amidotransferase content in the plant.

- Depending on the choice of the promoter, expression may take
- 20 place specifically in the leaves, in the seeds or in other parts of the plant. Such transgenic plants and their propagation material and their plant cells, tissue or parts are another subject of the present invention.

- 25 The invention will now be illustrated by the examples which follow, without being limited thereto.

## Examples

- 30 Recombinant methods on which the use examples are based:

## General cloning methods

- Cloning methods such as restriction cleavages, agarose gel
- 35 electrophoresis, purification of DNA fragments, transfer of nucleic acids to nitrocellulose and nylon membranes, linking DNA fragments, transformation of *Escherichia coli* cells, growing bacteria and the sequence analysis of recombinant DNA were carried out as described by Sambrook et al. (1989) (Cold Spring
- 40 Harbor Laboratory Press: ISBN 0-87969-309-6).

## Sequence analysis of recombinant DNA

- Recombinant DNA molecules were sequenced using an ABI laser
- 45 fluorescence DNA sequencer, following the method of Sanger (Sanger et al., Proc. Natl. Acad. Sci. USA, 74(1977), 5463-5467). Fragments resulting from a polymerase chain reaction were

## 15

sequenced and checked to avoid polymerase errors in constructs to be expressed.

## Analysis of total RNA from plant tissues

5 Total RNA from plant tissues was isolated as described by Logemann et al. (Anal. Biochem. 163(1987), 21). For the analysis, in each case 20 µg of RNA were separated in a formaldehyde-containing 1.5% agarose gel and transferred to nylon  
10 membranes (Hybond, Amersham). Specific transcripts were detected as described by Amasino (Anal. Biochem. 152(1986), 304). The DNA fragments employed as probe were radiolabeled with a Random Primed DNA Labeling Kit (Roche, Mannheim) and hybridized by standard methods (see Hybond instructions, Amersham).  
15 Hybridization signals were visualized by autoradiography with the aid of Kodak X-OMAT AR films.

Unless otherwise specified, the chemicals used were analytical grade and obtained from Fluka (Neu-Ulm), Merck (Darmstadt), Roth (Karlsruhe), Serva (Heidelberg) and Sigma (Deisenhofen). Solutions were made with a refined, pyrogen-free water, termed H<sub>2</sub>O hereinbelow, from a Milli-Q water refining system (Millipore, Eschborn). Restriction endonucleases, DNA-modifying enzymes and molecular biologic kits were obtained from AGS (Heidelberg), Amersham (Braunschweig), Biometra (Göttingen), Roche (Mannheim), Genomed (Bad Oeynhausen), New England Biolabs (Schwalbach/Taunus), Novagen (Madison, Wisconsin, USA), Perkin-Elmer (Weiterstadt), Pharmacia (Freiburg), Qiagen (Hilden) and Stratagene (Heidelberg). Unless otherwise specified, they were used in accordance with the manufacturer's instructions.

The bacterial strains used hereinbelow (*E. coli*, XL-1 Blue) were obtained from Stratagene. *E. coli* AT 2465 was obtained from the *coli* genetic stock center (Yale University, New Haven). The  
35 agrobacterial strain used for transforming plants (*Agrobacterium tumefaciens*, C58C1 with plasmid pGV2260 or pGV3850kan) was described by Deblaere et al. (*Nucl. Acids Res.* 13 (1985), 4777). Alternatively, it is also possible to use the agrobacterial strain LBA4404 (Clontech) or other suitable strains. Vectors  
40 which can be used for cloning are pUC19 (Yanish-Perron, *Gene* 33(1985), 103-119), pBluescript SK- (Stratagene), pGEM-T (Promega), pZerO (Invitrogen), pBin19 (Bevan et al., *Nucl. Acids Res.* 12(1984), 8711-8720) and pBinAR (Höfgen and Willmitzer, *Plant Science* 66 (1990), 221-230).

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0050/50796

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Following restriction and sequence analysis, two different clones were identified, Ntpur1.1 (clone 7.2) containing the DNA sequence SEQ-ID No. 1 and Ntpur1.2 (clone 9.2) containing the DNA sequence SEQ-ID No. 3, which encode reading frames with homology to *Arabidopsis thaliana* AtATase1. The amino acid sequences of Ntpur1.1 (SEQ-ID No. 2 - length: 573 amino acids) and Ntpur1.2 (SEQ-ID No. 4 - length: 573 amino acids) show 97% identity, see Table 1. The homology with AtATase1 at amino acid level is 81% in the case of Ntpur1.1 and 85% in the case of Ntpur1.2. The continuous reading frames start with nucleotide base 49 (Ntpur1.1) and 25 (Ntpur1.2) respectively, and are translated into polypeptides 573 amino acids in length.

Table 1

Amino acid comparison Ntpur1.1 x Ntpur1.2:

	1	MAATVSTASAAATNKSPLSQPLDKPFCSPSQKLLSLSPKTLPKPYRTLVT	50
	1	MAATVSTASAAATNKYPLSQPLDKPFCSLSQKLLSLSPKTHPKPYRTLIT	50
20	51	ASSKNPLNDVVSFKKSADNTLDSYFDDDDKPREECGVVGIIYGDSEASRLC	100
	51	ASSKNPLNDVISFKKSADNTLDSYFDDDDKPREECGVVGIIYGDSEASRLC	100
	101	YLALHALLHRGQEGAGIVAVNDDVLKSITGVGLVSDVFNESKLDQLPGDM	150
25	101	YLALHALQHRGQEGAGIVAVNDDVLKSITGVGLVSDVFNESKLDQLPGDM	150
	151	AIGHVWYSTAGSSMLKNVQPFVANYKFGSVGVAHNGNLVNYKLLRGELEE	200
	151	AIGHVRYSTAGSSMLKNVQPFVASYKFGSVGVAHNGNLVNYKLLRSELEE	200
30	201	NGSIFNTSSDTEVVLHLIAISKARPFLLRIVEACEKIEGAYSMVFVTEDEK	250
	201	NGSIFNTSSDTEVVLHLIAISKARPFLLRIVEACEKIEGAYSMVFVTEDEK	250
	251	LVAVRDPHGFRPLVMGRRSNGAVVFASSETCALDLIEATYEREVNPGEVVV	300
	251	LVAVRDPHGFRPLVMGRRSNGAVVFASSETCALDLIEATYEREVNPGEVVV	300
35	301	VDKDGVHSHIYLMPPHPEHKSCIFEHIYFALPNSVVFGRSVYESRRAFGEIL	350
	301	VDKDGVQSICLMPHPERKSCIFEHIYFALPNSVVFGRSVYESRRAFGEIL	350
	351	ATEAPVECDVGIAVPDPSGIVAALGYAAKAGVFPQQGLIRSHYVGRFTFIEP	400
40	351	ATEAPVECDVVIAPDPSGVVAALGYAAKAGVFPQQGLIRSHYVGRFTFIEP	400
	401	SQKIRDFGVKLKLSPVRALEGGKRVVVVDDSIVRGTTSSKIVRLLKEAGA	450
	401	SQKIRDFGVKLKLSPVRAVLEGGKRVVVVDDSIVRGTTSSKIVRLLKEAGA	450
45	451	KEVHMRIASPPPIIASCYYGVDTPSSDELISNRMSVEEIKEFIGSDSLAFL	500
	451	KEVHMRIASPPPIIASCYYGVDTPSSDELISNRMSVEEIKEFIGSDSLAFL	500

0050/50796

18

501 PMDSLNLKLLGNDKSFYACFSGNYVPVEPTGKVKRIGDFMDDGLSGDMDS 550

|||||

501 PMDSLNLKLLGNDKSFYACFSGNYVPVEPTGKVKRIGDFMDDGLSGDMDS 550

551 IDGGWLPGSSRVQKILNEVRTG 573

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551 IDGGWLPGSSRVQKILNEVRTS 573

Compared with bacterial and human PRPP amidotransferase sequences, the plant proteins (Ntpur1.1, Ntpur1.2, AtATase1) show an extended N-terminus with a large proportion of basic amino acids (Table 2), which suggests the function of a transit peptide for plastid import (von Heijne et al., Eur. J. Biochem. 180(1989), 535-545).

Table 2

Sequence comparison of Arabidopsis thaliana (AtATase1), Bacillus subtilis (BacSu\_purF), Human (pur1\_hum) and Nicotiana tabacum (Ntpur1.1), Ntpur1.2) PRPP amidotransferase proteins.

		1				50
20	AtATase1	~~~~~	~~~~~	~~~~~SLN	QTILLTPINL	SLSSPNPSLN
	BacSu_purF	~~~~~	~~~~~	~~~~~	~~~~~	~~~~~
	Ntpur1	LAPHLFLLS	SFFPPPMAAT	VSTASAAATN	KSPLSQPLDK	PFCSPSQKL.
	Ntpur1-2	~~~~~LS	SFFPPPMAAT	VSTASAAATN	KYPLSQPLDK	PFCSLSQKL.
	pur1_hum	~~~~~	~~~~~	~~~~~	~~~~~	~~~~~
		51				100
25	AtATase1	LHISLS.FLL	PSPLLLLHSS	MESPPTSPLL	HHPKNNSHAP	FDYHNDEDDE
	BacSu_purF	~~~~~	~~~~~	~~~~~	~~~~~MLAEIK	~~~~~
	Ntpur1	..LSLSPKTL	PKPYRTLVT	SSKNPLNDVV	SFKKSADNTL	DSYFDDDED..
	Ntpur1-2	..LSLSPKTH	PKPYRTLIT	SSKNPLNDVI	SFKKSADNTL	DSYFDDDDD..
	pur1_hum	~~~~~	~~~~~	~~~~~	~~~~~	~~~~~MELEEL
		101				150
30	AtATase1	KPREECGVVG	IYGDPE....	..ASRLFYLA	LHALQHRGQE	GAGIVTVSPE
	BacSu_purF	GLNEECGVFG	IWGHEE....	..APQITYYG	LHSLQHRGQE	GAGIVATDGE
	Ntpur1	KPREECGVVG	IYGDSE....	..ASRLCYLA	LHALLHRGQE	GAGIVAVN.D
	Ntpur1-2	KPREECGVVG	IYGDSE....	..ASRLCYLA	LHALQHRGQE	GAGIVAVN.D
	pur1_hum	GIREECGVFG	CIASGEWPTQ	LDVPHVITLG	LVGLQHRGQE	SAGIVTSDGS
35		151				200
	AtATase1	KV..LQTITG	VGLVSEVFNE	SKLDQL.PGE	FAIAHVRYST	AGASMLKNVQ
	BacSu_purF	K...LTAHKG	QGLITEVFQN	GELSKV.KGK	GAIGHVRYAT	AGGGGYENVQ
	Ntpur1	DV..LKSITG	VGLVSDVFNE	SKLDQL.PGD	MAIGHVWYST	AGSSMLKNVQ
	Ntpur1-2	DV..LKSITG	VGLVSDVFNE	SKLDQL.PGD	MAIGHVRYST	AGSSMLKNVQ
40	pur1_hum	SVPTFKSHKG	MGLVNHVFTE	DNLKKLYVSN	LGIGHTRYAT	TGKCELENCQ
		201				250
	AtATase1	PFV.AGYRFG	SIGVAHNGNL	VNYKTLRAML	EENGSIFFNTS	SDTEVVLHLI
	BacSu_purF	PLLFRSQNNG	SLALAHNGNL	VNATQLKQQL	ENQGSIFQTS	SDTEVLAHLI
	Ntpur1	PFV.ANYKFG	SVGVAHNGNL	VNYKLLRGEL	EENGSIFFNTS	SDTEVVLHLI
	Ntpur1-2	PFV.ASYKFG	SVGVAHNGNL	VNYKLLRSEL	EENGSIFFNTS	SDTEVVLHLI
45	pur1_hum	PFVVETLH.G	KIAVAHNGEL	VNAARLRKKL	LRHGIGLSTS	SDSEMITQLL
		251				300
	AtATase1	AISKAR....	..PFFMRIID	ACEKLQGAYS	MVFVTEKLV	AVRDPYGFRR



0050/50796

## 19

	BacSu_purF	KRSGHF....	..TLKDQIKN	SLSMLKGAYA	FLIMTETEMI	VALDPNGLRP
	Ntpur1	AISKAR....	..PFLLRIVE	ACEKIEGAYS	MVFVTEDEKL	AVRDPHGFRP
	Ntpur1-2	AISKAR....	..PFLLRIVE	ACEKIEGAYS	MVFVTEDEKL	AVRDPHGFRP
	pur1_hum	AYTPPQEQDD	TPDWVARIKN	LMKEAPTAYS	LLIMHRDVIY	AVRDPYGNRP
		301				350
5	AtATasel	LVMGR.....	.....R	SNGAVVFASE	TCALDLIEAT	YEREVYPGEV
	BacSu_purF	LSIGM.....	.....M	GD.AYVVASE	TCAFDVVGAT	YLREVEPGEM
	Ntpur1	LVMGR.....	.....R	SNGAVVFASE	TCALDLIEAT	YEREVNPGEV
	Ntpur1-2	LVMGR.....	.....R	SNGAVVFASE	TCALDLIEAT	YEREVNPGEV
	pur1_hum	LCIGRLIPVS	DINDKEKSTS	ETEGWVVSSE	SCSFLSIGAR	YYREVLPGEI
		351				400
10	AtATasel	LVVDKDGVS	QCLMPKFEPK	Q...CIFEHI	YFSLPNSIVF	GRSVYESRHH
	BacSu_purF	LIINDEGMKS	ERFSMNINRS	I...CSMEYI	YFSRPDSNID	GINVHSARKN
	Ntpur1	VVVDKDGVS	IYLMPPHEHK	S...CIFEHI	YFALPNSVVF	GRSVYESRRA
	Ntpur1-2	VVVDKDGVS	ICLMPHPERK	S...CIFEHI	YFALPNSVVF	GRSVYESRRA
	pur1_hum	VEISRHNVT	LDIISRSEGN	PVAFCIFEYV	YFARPDMSFE	DQMVTVRYR
		401				450
15	AtATasel	FGEILATESP	VECDVVIAPV	DSGVVAALGY	AAKSGVPFQQ	GLIRSHYVGR
	BacSu_purF	LGMMLAQESA	VEADVVTGVP	DSSISAAIGY	AEATGIPYEL	GLIKNRYVGR
	Ntpur1	FGEILATEAP	VECDVGIAPV	DSGIVAALGY	AAKAGVPFQQ	GLIRSHYVGR
	Ntpur1-2	FGEILATEAP	VECDVVIAPV	DSGVVAALGY	AAKAGVPFQQ	GLIRSHYVGR
20	pur1_hum	CGQQLAIEAP	VDADLVSTVP	ESATPAALAY	AGKCGLPYVE	VLCKNRYVGR
		451				500
25	AtATasel	TFIEPSQKIR	DFGVKLLKSP	VRGVLEGKRV	VVVDDSIVRG	TTSSKIVRLL
	BacSu_purF	TFIQPSQALR	EQGVRMKLSA	VRGVVEGKRV	VMVDDSIVRG	TTSSRIVTML
	Ntpur1	TFIEPSQKIR	DFGVKLLKSP	VRALLEGKRV	VVVDDSIVRG	TTSSKIVRLL
	Ntpur1-2	TFIEPSQKIR	DFGVKLLKSP	VRVLEGGKRV	VVVDDSIVRG	TTSSKIVRLL
	pur1_hum	TFIQPNMRLR	QLGVAKKFGV	LSDNFKGKRI	VLVDDSIVRG	NTISPIIKLL
		501				550
30	AtATasel	REAGAKEVHM	RIASPPIVAS	CYYGVDTPSS	EELISNRLSV	EEINEFIGSD
	BacSu_purF	REAGATEVHV	KISSPPIAHP	CFYGIDTSTH	EELIASSHSV	GEIRQEIGAD
	Ntpur1	KEAGAKEVHM	RIASPPIIAS	CYYGVDTPSS	DELISNRMSV	EEIKEFIGSD
	Ntpur1-2	KEAGAKEVHM	RIASPPIIAS	CYYGVDTPSS	DELISNRMSV	EEIKEFIGSD
	pur1_hum	KESGAKEVHI	RVASPPIKYP	CFMGINIPTK	EELIANKPEF	DHLAEYLGAN
		551				600
35	AtATasel	SLAFLSFDTL	KKHL.....	.....GK...	.DSK.SFCYA	
	BacSu_purF	TLSFLSVEGL	LKGI.....	.....GRKYD	.DSNCGQCLA	
	Ntpur1	SLAFLPMDSL	NKLL.....	.....GN...	.DSK.SFCYA	
	Ntpur1-2	SLAFLPMDSL	NKLL.....	.....GN...	.DSK.SFCYA	
	pur1_hum	SVVYLSVEGL	VSSVQEGIKF	KKQKEKKHDI	MIQENGNGLE	CFEKSGHCTA
		601				650
40	AtATasel	CFTGDYPVKP	TEVKVKRGGG	DFIDDGLVGS	FENIEAGWVR	-----
	BacSu_purF	CFTGKYPTI	YQDVTLPVK	EAVLTK----	-----	-----
	Ntpur1	CFSGNYPVEP	TG.KVKR.IG	DFMDDGLSGD	MDSIDGGWLP	GSSRVQKTI
	Ntpur1-2	CFSGNYPVEP	TG.KVKR.IG	DFMDDGLSGD	MDSIDGGWLP	GSSRVQKTI
	pur1_hum	CLTGKYPVEL	EW-----	-----	-----	-----
		651				
45	AtATasel	-----				
	BacSu_purF	-----				
	Ntpur1	NEVRTG				
	Ntpur1-2	NEVRTS				
	pur1_hum	-----				

0050/50796

## 20

### Example 2

#### Expression of tobacco PRPP amidotransferase in E. coli

- 5 The purpose of expressing Ntpur1.2 in E. coli was to prove that the Ntpur1.2-encoded PRPP amidotransferase enzyme was active. To this end, a 1523 bp fragment was amplified in a PCR with Pfu polymerase using the oligonucleotides Jle336:  
5'-ttttgctagcgcactcgtattttgacg-3' and Jle337:  
5'-aaaaagatctcaggttctaacttcat -3' and Ntpur1.2 DNA as template.
- 10 The DNA fragment generated encodes a PRPP amidotransferase enzyme with is truncated N-terminally by 86 amino acids and no longer contains the transit peptide to be received. This truncated form of PRPP amidotransferase enzyme starts N-terminally with the
- 15 amino acids MDSYFDDDD. Using the oligonucleotides, an NheI cleavage site and a BglII cleavage site were inserted via which the fragment generated was ligated into the NheI- and BamHI-cleaved expression vector pET11a (Novagen).
- 20 For expression, the E. coli strain BL21(DE3)LysS (Novagen) was transformed with the construct pETNtpur1.2 which had thus been generated. Following overnight culture, a day culture was inoculated to OD<sub>600</sub> = 0.1 and, after an OD<sub>600</sub> = 0.7 had been reached, induced with 1mM IPTG. A total cell extract was produced
- 25 by the pressure disruption method ("French press") in 50mM Tris-HCl, pH 7.4; 150mM NaCl. Following SDS polyacrylamide gel electrophoresis, an overexpressed protein of approx. 65 kDa was excised from the gel. To produce antisera, the protein was injected into rabbits (contractor: Eurogentec, Herstal, Belgium).

30

### Example 3

#### Assay system for measuring the activity of plant PRPP amidotransferase activity

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- The above-described method for measuring plant PRPP amidotransferase activity by the method of Reynolds et al. (Archives of Biochemistry and Biophysics 229 (1984), 623-631) is not suitable for high-throughput assaying owing to the use of
- 40 radioactive materials. This is why an alternative assay system with which the plant PRPP amidotransferase activity is detected in the protein extract is detected on the basis of the formation of the reaction product glutamate, based on the method described by Shid and Ishii (Journal of Biological Chemistry 66 (1969),
- 45 175-181) for E. coli PRPP amidotransferase. The concentration of

0050/50796

## 21

the glutamate which forms is measured by converting it with glutamate dehydrogenase (GluDH) and monitoring APADH formation photometrically at 363 nm.



- 10 (PRPP = phosphoribosyl pyrophosphate, PRA = phosphoribosylamine, APAD = 3-acetylpyridineadenin dinucleotide, PRAT = PRPP amidotransferase)

To this end, the reaction batch (see below) is incubated at 37°C

- 15 for up to 60 minutes and the reaction was quenched by incubation at 95°C for 5 minutes.

Reaction batch:

20	375 $\mu$ l	100 mM	Tris/HCl buffer pH 8.0
	75 $\mu$ l	100 mM	MgCl <sub>2</sub>
	75 $\mu$ l	30 mM	phosphoribosyl pyrophosphate
	75 $\mu$ l	100 mM	L-glutamine
	50 $\mu$ l		H <sub>2</sub> O
25	<u>100 <math>\mu</math>l</u>		protein extract
	750 $\mu$ l		

The glutamate formed was detected in the detection batch (see below) by measuring the increase in APADH photometrically at

- 30 363 nm following addition of glutamate dehydrogenase.

Detection batch:

	375 $\mu$ l	100 mM	Tris/HCl buffer pH 8.0
35	75 $\mu$ l	500 mM	KCl
	125 $\mu$ l		H <sub>2</sub> O
	75 $\mu$ l	3 mM	APAD
	<u>100 <math>\mu</math>l</u>		of the reaction batch
	750 $\mu$ l		

40

Start of the detection reaction with 2  $\mu$ l (approx. 4 units) glutamate dehydrogenase (Sigma).

The assay system lends itself in particular for measuring PRPP

- 45 amidotransferase activity from plant material and in expression extracts, for example from baculovirus-infected insect cells.

## 22

## Example 4

Functional expression of tobacco PRPP amidotransferase in insect cells

5

The back-to-back expression system from GibcoBRL was employed for expressing Ntpur1.1 in baculovirus-infected insect cells. To this end, Ntpur1.1 was employed in a PCR. The reaction mixtures contained approximately 1 ng/ $\mu$ l Ntpur1.1 DNA, 0.5  $\mu$ M of the

10 oligonucleotides 5'-tat agg atc cat gga ctc cta ttt tga cg-3' and 5'-atg aat tct agc tgg ttc taa ctt c-3', 200  $\mu$ M deoxynucleotides (Pharmacia), 0.04 U/ $\mu$ l Pfu polymerase (Stratagene) and buffer conditions were set following the manufacturer's instructions.

15 The amplification conditions were set as follows:

## Step 1:

Denaturation temperature: 95°C, 0.5 min  
20 Annealing temperature: 40°C, 0.5 min  
Elongation temperature: 72°C, 2 min  
Number of cycles for Step 1: 2

## Step 2:

25

Denaturation temperature: 95°C, 0.5 min  
Annealing temperature: 50°C, 0.5 min  
Elongation temperature: 72°C, 3 min  
Number of cycles for Step 2: 25

30

The PCR product was ligated into the StuI-cut vector pFastBac1 (GibcoBRL). The correct orientation of the insert was ensured by control digest with KpnI. The resulting transfer vector pFastBacNtpur1.2 was used following the manufacturer's

35 instructions for generating recombinant baculoviruses by means of Sf21 insect cells (Invitrogen). Sf21 insect cells were infected with the recombinant baculovirus (BvNtpur1.2). After 2-4 days, the cells were harvested by centrifugation. A protein of approx. 54kDa, which corresponds to the expected size of PRPP

40 amidotransferase, was identified in the total extract by SDS polyacrylamide gel electrophoresis. A total cell extract was prepared by the pressure disruption method ("French press") in extraction buffer (100 mM HEPES pH 8.0; 2.5 mM EDTA; 10% glycerol; 20 mM DTE; 0.2 mM PEFA block) and, after being freed  
45 from salt over a PD10 column (Pharmacia), used for measuring PRPP amidotransferase activity in the assay described (see Example 3).

0050/50796

23

Example 5

Generation of plant transformation vectors

- 5 To generate binary vectors for plant transformation, clone Ntpur1.1 was cleaved with SmaI and EcoRV, and a fragment comprising 1482 bp was isolated and ligated into the SmaI-cleaved vector pBinAR (Höfgen and Willmitzer, Plant Science 66(1990), 221-230). The antisense and sense constructs thus obtained were  
10 termed pBinAR-Ntpur1A and pBinAR-Ntpur1, respectively; see Figure 1.

Example 6

15 Generation of transgenic tobacco plants

- Plasmid pBinAR-Ntpur1A and pBinAR-Ntpur1 were transformed into Agrobacterium tumefaciens C58C1:pGV2260 (Deblaere et al., Nucl. Acids. Res. 13(1984), 4777-4788). To transform tobacco plants  
20 (Nicotiana tabacum cv. Samsun NN), a 1:50 dilution of an overnight culture of a positively transformed agrobacterial colony in Murashige-Skoog medium (Murashige and Skoog, Physiol. Plant. 15(1962), 473) supplemented with 2% sucrose (2MS medium) was used. Leaf disks of sterile plants (in each case approx.  
25 1 cm<sup>2</sup>) were incubated in a Petri dish for 5-10 minutes with a 1:50 agrobacterial dilution. This was followed by 2 days' incubation in the dark at 25°C on 2MS medium with 0.8% Bacto agar. Cultivation was continued after 2 days and at 16 hours light/8 hours dark and continued in a weekly rhythm on MS medium  
30 with 500 mg/l claforan (cefotaxime-sodium), 50 mg/l kanamycin, 1 mg/l benzylaminopurine (BAP), 0.2 mg/l naphthylacetic acid and 1.6 g/l glucose. Growing shoots were transferred to MS medium supplemented with 2% sucrose, 250 mg/l claforan and 0.8% Bacto agar.  
35 Regenerated shoots were obtained on 2MS medium with kanamycin and claforan, transferred into soil after rooting, and, after cultivation for two weeks in a controlled-environment cabinet in a 16-hour light/8-hour dark rhythm at 60% atmospheric humidity,  
40 analyzed for PRPP amidotransferase expression and activity and for altered metabolite contents and phenotypic growth characteristics. Altered nucleotide contents can be determined for example following the method of von Stitt et al., FEBS Letters 145(1982), 217-222.

0050/50796

## 24

## Example 7

## Analysis of transgenic plants

5 Transgenic plants which were transformed with the construct with pBinAR-Ntpurl are characterized by a growth which is reduced by different degrees and by large-scale bleaching of the leaves in comparison with untransformed control plants (Fig. 2). RNA analysis by the Northern blot technique showed a reduced amount  
10 of Ntpurl.1-RNA in transgenic lines with the above-described phenotype (Fig. 3). These effects were also observed in subsequent generations of the transgenic lines.

To test the correlation with growth reduction, PRPP  
15 amidotransferase activity in the transgenic lines was measured and compared with that in untransformed controls. To this end, in each case approx. 30 g of leaves from plants approximately 20 cm in height were homogenized with 50 ml of extraction buffer at +4°C.

20

## Extraction buffer:

100 mM	HEPES pH 8,0
2.5 mM	EDTA
25 10%	glycerol
20 mM	DTE
0,2 mM	PEFA block (40mM)

The disruption extract was filtered through Miracloth  
30 (Calbiochem, Bad Soden) and spun at 16,000 rpm in a Sorval centrifuge. The resulting supernatant was precipitated with ammonium sulfate at 4°C. The 30% - 60% fraction was solubilized in an extraction buffer and freed from salt by means of a PD-10 column (Pharmacia, Sweden). The extract thus obtained is stable  
35 for at least 24 hours and can be stored over a prolonged period at -20°C after addition of glycerol (end concentration 50%). The extract can be employed directly in the activity determination. Compared to wild-type plants, the PRPP amidotransferase activity in the transgenic lines was markedly reduced, see Fig. 4. Fig. 4A  
40 shows the PRPP amidotransferase activity based on the protein quantity. Fig. 4B shows the PRPP amidotransferase activity based on the fresh weight.

0050/50796

## 25

These data establish a direct connection between reduced PRPP amidotransferase activity and reduced growth of the tobacco plants and thus identify PRPP amidotransferase for the first time as suitable target protein for herbicidal active ingredients.

5

### Example 8

Search for PRPP amidotransferase activity inhibitors

- 10 The in-vitro assay described in Example 3 can be used together with high-throughput methods for searching for PRPP amidotransferase activity inhibitors. To this end, the PRPP amidotransferase activity can be prepared from plant tissue, see Example 7. Alternatively, a plant PRPP amidotransferase can be
- 15 expressed in *E. coli*, insect cells or in another suitable expression system. Known PRPP amidotransferase inhibitors such as glutamine antagonists were identified in this manner.

### Example 9

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Analysis of the adenine and guanine nucleotide contents in transgenic plants.

- Leaf material (in each case 5 disks of 6 mm diameter) was
- 25 harvested from wild-type plants and transgenic plants transformed with the construct pBinAR-Ntpurl and the subsequent generation (lines 3.1, 3.2, 3.9, 25.1 and 38.8) and frozen immediately in liquid nitrogen. TCA extracts were subsequently prepared by standard methods and employed for the determination of the
- 30 nucleotide contents.

- In the transgenic plants, with the exception of line 38.8, AMP is reduced greatly in the green regions of the leaf and less in the yellow regions of the leaf compared to the wild type (WT) (see
- 35 Fig. 5).

No changes compared to the wild type were observed for the guanosine nucleotide GTP, GDP and GMP.

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We claim:

1. A DNA sequence containing the encoding region of a plant PRPP  
amidotransferase, wherein this DNA sequence has the  
nucleotide sequence SEQ-ID No. 1 or SEQ-ID No. 3.
2. A DNA sequence hybridizing with the DNA sequence SEQ-ID No. 1  
or SEQ-ID No. 3 as claimed in claim 1 or parts thereof or  
derivatives, derived from this sequence by insertion,  
deletion or substitution and encoding a protein which has the  
biological activity of a PRPP amidotransferase.
3. A protein with PRPP amidotransferase activity comprising an  
amino acid sequence which constitutes a subsequence of at  
least 100 amino acids from SEQ-ID No. 2 or SEQ-ID No. 4.
4. A protein as claimed in claim 3, which comprises, as amino  
acid sequence, the subsequence 100 - 450 from SEQ-ID No. 2 or  
SEQ-ID No. 4.
5. A protein as claimed in claim 4, which comprises, as amino  
acid sequence, the sequence shown in SEQ-ID No. 2 or SEQ-ID  
No. 4.
6. The use of a DNA sequence as claimed in claim 1 or 2 for  
introduction into pro- or eukaryotic cells, this sequence  
optionally being linked to control elements which ensure  
transcription and translation in the cells and leading to the  
expression of a translatable mRNA which causes the synthesis  
of a plant PRPP amidotransferase.
7. The use of a DNA sequence as claimed in claim 1 or 2 for  
generating an assay system for identifying herbicidally  
active plant PRPP amidotransferase inhibitors.
8. A method of finding herbicidally active substances which  
inhibit the activity of the plant PRPP amidotransferase,  
which comprises preparing, in a first step, PRPP  
amidotransferase using a DNA sequence as claimed in claim 1  
or 2 and measuring, in a second step, the activity of the  
plant PRPP amidotransferase in the presence of a test  
substance.
9. The method as claimed in claim 9, wherein the plant PRPP  
amidotransferase is measured in a high-throughput screening  
(HTS). The method as claimed in claim 9, wherein the plant



0050/50796

27

PRPP amidotransferase is measured in a high-throughput screening (HTS).

10. A method of identifying herbicidally active substances which inhibit the PRPP amidotransferase activity in plants, with the following steps:
- a) the generation of transgenic plants, plant tissues or plant cells which comprise an additional DNA sequence encoding an enzyme with PRPP amidotransferase activity and which are capable of overexpressing an enzymatically active PRPP amidotransferase;
  - b) applying a substance to transgenic plants, plant cells, plant tissue or plant parts and to untransformed plants, plant cells, plant tissue or plant parts;
  - c) determining the growth or the viability of the transgenic and the untransformed plants, plant cells, plant tissue or plant parts after application of the chemical substance; and
  - d) comparing the growth or the viability of the transgenic and the untransformed plants, plant cells, plant tissue or plant parts after applying the chemical substance;
- where a suppression of the growth or the viability of the untransformed plants, plant cells, plant tissue or plant parts, but an absence of potent suppression of the growth or viability of the transgenic plants, plant cells, plant tissue or plant parts, confirms that the substance of b) shows herbicidal activity and inhibits the PRPP amidotransferase enzyme activity in plants.
11. An assay system based on the expression of a DNA sequence SEQ-ID No. 1 or SEQ-ID No.9 as claimed in claim 1 or 2 for identifying herbicidally active plant PRPP amidotransferase inhibitors.
12. An assay system as claimed in claim 11 for identifying herbicidally active plant PRPP amidotransferase inhibitors, which comprises incubating the enzyme with a test substrate to be studied and, after a suitable reaction time, determining the enzymatic activity of the enzyme in comparison with the activity of the uninhibited enzyme.

0050/50796

## 28

13. A plant PRPP amidotransferase inhibitor.

14. A plant PRPP amidotransferase inhibitor identified using an assay system as claimed in claim 11 or 12.

5

15. An inhibitor as claimed in claim 13 or 14 for use as herbicide.

10 16. A method of eliminating undesired vegetation, which comprises treating the plants to be eliminated with a compound which binds specifically to PRPP amidotransferase encoded by a DNA sequence as claimed in claim 1 or 2 and which inhibits its function.

15 17. A plant with a modified purine nucleotide content, generated by additionally expressing a DNA sequence SEQ-ID No. 1 or SEQ-ID No. 3 as claimed in claim 1 or 2 in sense or antisense orientation.

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0050/50796

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## PRPP amidotransferase

## Abstract

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The present invention relates to DNA sequences encoding a polypeptide with PRPP amidotransferase (EC 2.4.2.14) activity. The invention furthermore relates to the use of these nucleic acids for generating a test system.

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The diagram illustrates the Nt-pur1.1 gene construct. It begins with a 35S promoter (represented by a trapezoid) followed by an EcoRI site (indicated by an upward arrow). The Nt-pur1.1 gene (1482 bp) is shown as a rectangle with an internal arrow indicating the direction of transcription. A SmaI site is located at the end of the 35S promoter (indicated by a downward arrow). The gene is flanked by an OCS terminator (represented by a rectangle). A SmaI site is also located at the end of the gene (indicated by a downward arrow). The distance between the two SmaI sites is labeled as „EcoRV-SmaI“.

Fig. 2

2/4

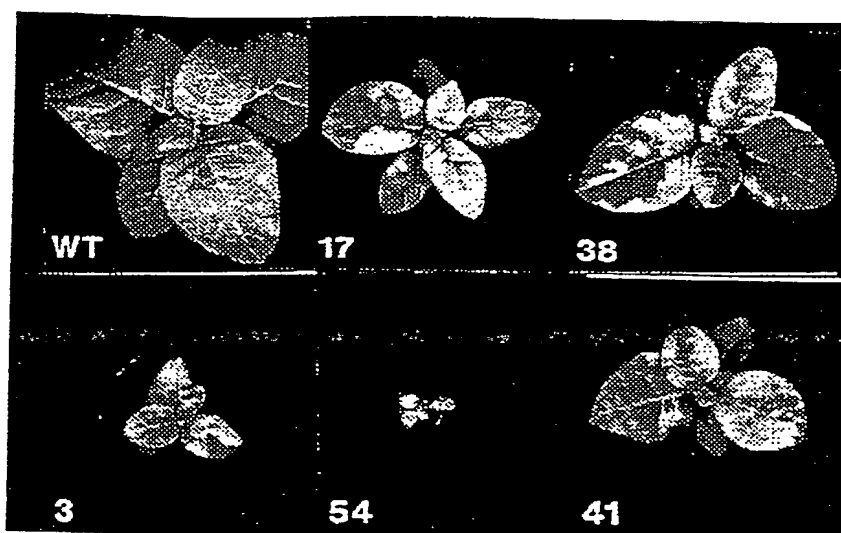


Fig. 3

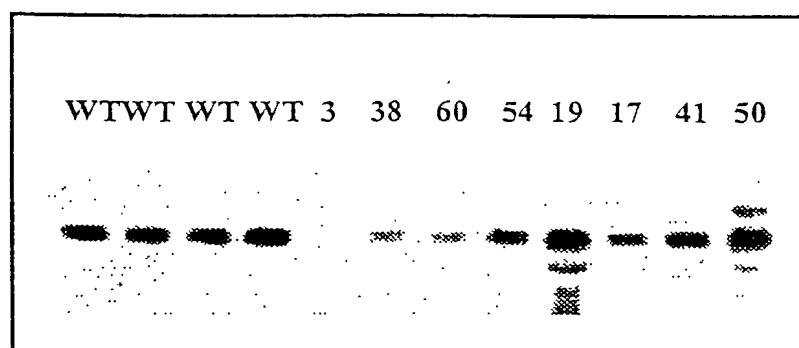


Fig. 4

3/4

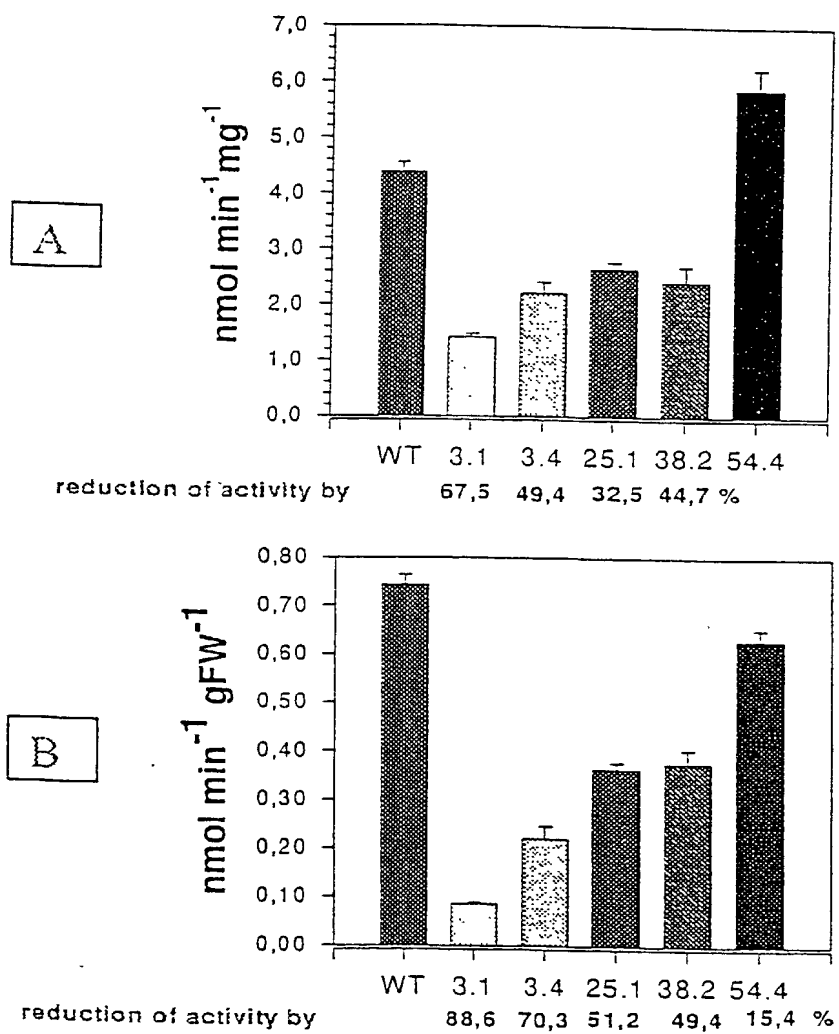
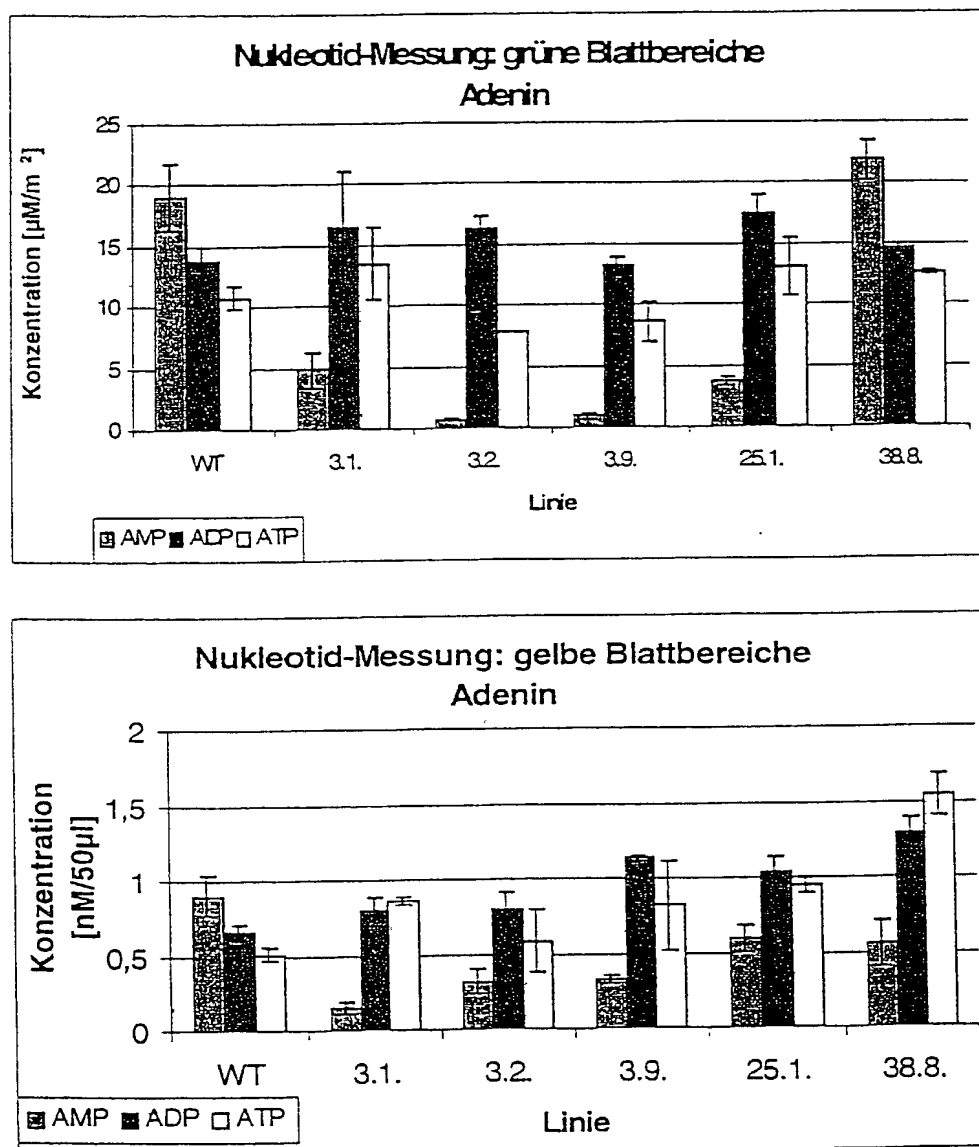


Fig. 5



# Declaration, Power of Attorney and Petition

Page 1 of 3

0050/050796

Customer No.

We (I), the undersigned inventor(s), hereby declare(s) that:

My residence, post office address and citizenship are as stated below next to my name,

We (I) believe that we are (I am) the original, first, and joint (sole) inventor(s) of the subject matter which is claimed and for which a patent is sought on the invention entitled

PLANT PRPP AMIDOTRANSFERASE

the specification of which

☐ is attached hereto.

☐ was filed on \_\_\_\_\_ as

Application Serial No. \_\_\_\_\_

and amended on \_\_\_\_\_.

☒ was filed as PCT international application

Number PCT/EP/00/09839 \_\_\_\_\_

on 07 October 2000 \_\_\_\_\_,

and was amended under PCT Article 19

on \_\_\_\_\_ (if applicable).

We (I) hereby state that we (I) have reviewed and understand the contents of the above-identified specification, including the claims, as amended by any amendment referred to above.

We (I) acknowledge the duty to disclose information known to be material to the patentability of this application as defined in Section 1.56 of Title 37 Code of Federal Regulations.

We (I) hereby claim foreign priority benefits under 35 U.S.C. § 119(a)-(d) or § 365(b) of any foreign application(s) for patent or inventor's certificate, or § 365(a) of any PCT International application which designated at least one country other than the United States, listed below and have also identified below, by checking the box, any foreign application for patent or inventor's certificate, or PCT International application having a filing date before that of the application on which priority is claimed. Prior Foreign Application(s)

Application No.	Country	Day/Month/Year	Priority Claimed
19949000.7	Germany	11 October 1999	<input checked="" type="checkbox"/> Yes <input type="checkbox"/> No



We (I) hereby claim the benefit under Title 35, United States Codes, § 119(c) of any United States provisional application(s) listed below.

_____	_____
(Application Number)	(Filing Date)
_____	_____
(Application Number)	(Filing Date)

We (I) hereby claim the benefit under 35 U.S.C. § 120 of any United States application(s), or § 365(c) of any PCT International application designating the United States, listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States or PCT International application in the manner provided by the first paragraph of 35 U.S.C. § 112, I acknowledge the duty to disclose information which is material to patentability as defined in 37 CFR § 1.56 which became available between the filing date of the prior application and the national or PCT International filing date of this application.

Application Serial No.	Filing Date	Status (pending, patented, abandoned)
_____	_____	_____
_____	_____	_____
_____	_____	_____

And we (I) hereby appoint Nixon & Vanderhye P.C., Attorneys at Law, 1100 North Glebe Road, Arlington, Virginia 22201-4714, our attorneys, with full power of substitution and revocation, to prosecute this application, to make alterations and amendments therein, to sign the drawings, to receive the patent, and to transact all business in the Patent Office connected therewith.

We (I) declare that all statements made herein of our (my) own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issuing thereon.

**Declaration**

Page 3 of 3

0050/050796

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0050/50796

SEQUENCE LISTING

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0050/50796

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0050/50796

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0050/50796

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Ile	Phe	Glu	His	Ile	Tyr	Phe	Ala	Leu	Pro	Asn	Ser	Val	Val	Phe	Gly	
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Arg	Ser	Val	Tyr	Glu	Ser	Arg	Arg	Ala	Phe	Gly	Glu	Ile	Leu	Ala	Thr	
			340					345					350			
Glu	Ala	Pro	Val	Glu	Cys	Asp	Val	Gly	Ile	Ala	Val	Pro	Asp	Ser	Gly	
		355					360					365				
Ile	Val	Ala	Ala	Leu	Gly	Tyr	Ala	Ala	Lys	Ala	Gly	Val	Pro	Phe	Gln	
	370					375						380				
Gln	Gly	Leu	Ile	Arg	Ser	His	Tyr	Val	Gly	Arg	Thr	Phe	Ile	Glu	Pro	
385					390					395					400	
Ser	Gln	Lys	Ile	Arg	Asp	Phe	Gly	Val	Lys	Leu	Lys	Leu	Ser	Pro	Val	
				405					410					415		
Arg	Ala	Leu	Leu	Glu	Gly	Lys	Arg	Val	Val	Val	Val	Asp	Asp	Ser	Ile	
			420					425					430			
Val	Arg	Gly	Thr	Thr	Ser	Ser	Lys	Ile	Val	Arg	Leu	Leu	Lys	Glu	Ala	
		435					440					445				
Gly	Ala	Lys	Glu	Val	His	Met	Arg	Ile	Ala	Ser	Pro	Pro	Ile	Ile	Ala	
	450					455					460					
Ser	Cys	Tyr	Tyr	Gly	Val	Asp	Thr	Pro	Ser	Ser	Asp	Glu	Leu	Ile	Ser	
465					470					475					480	
Asn	Arg	Met	Ser	Val	Glu	Glu	Ile	Lys	Glu	Phe	Ile	Gly	Ser	Asp	Ser	
				485					490					495		
Leu	Ala	Phe	Leu	Pro	Met	Asp	Ser	Leu	Asn	Lys	Leu	Leu	Gly	Asn	Asp	
			500					505					510			
Ser	Lys	Ser	Phe	Cys	Tyr	Ala	Cys	Phe	Ser	Gly	Asn	Tyr	Pro	Val	Glu	
	515						520					525				
Pro	Thr	Gly	Lys	Val	Lys	Arg	Ile	Gly	Asp	Phe	Met	Asp	Asp	Gly	Leu	
	530					535					540					
Ser	Gly	Asp	Met	Asp	Ser	Ile	Asp	Gly	Gly	Trp	Leu	Pro	Gly	Ser	Ser	
545					550					555					560	



570

<213> Nicotiana tabacum

 $\langle 222 \rangle \quad (25) \dots (1743)$ 

Met Ala Ala Thr Val Ser Thr Ala Ser

5

25

40

55

70

85

105

120

135

ttc aat gag tca aag ctt gac caa ctc cct ggt gac atg gca att ggc	483
Phe Asn Glu Ser Lys Leu Asp Gln Leu Pro Gly Asp Met Ala Ile Gly	
140 145 150	
 cac gta agg tac tct act gct ggc tct tct atg tta aaa aat gtt cag	531
His Val Arg Tyr Ser Thr Ala Gly Ser Ser Met Leu Lys Asn Val Gln	
155 160 165	
 cct ttt gtt gct agt tat aaa ttt ggg tca gtt ggt gtt gcc cat aat	579
Pro Phe Val Ala Ser Tyr Lys Phe Gly Ser Val Gly Val Ala His Asn	
170 175 180 185	
 ggg aat tta gtg aat tat aag tta ctg cgt agt gaa cta gag gaa aat	627
Gly Asn Leu Val Asn Tyr Lys Leu Leu Arg Ser Glu Leu Glu Glu Asn	
190 195 200	
 ggg tca att ttt aat aca agt tct gat act gag gtt gta ctt cac ctt	675
Gly Ser Ile Phe Asn Thr Ser Ser Asp Thr Glu Val Val Leu His Leu	
205 210 215	
 att gct ata tct aaa gct agg cca ttt tta ttg agg att gtt gag gct	723
Ile Ala Ile Ser Lys Ala Arg Pro Phe Leu Leu Arg Ile Val Glu Ala	
220 225 230	
 tgt gaa aaa att gaa ggt gct tat tct atg gtg ttt gtt act gag gat	771
Cys Glu Lys Ile Glu Gly Ala Tyr Ser Met Val Phe Val Thr Glu Asp	
235 240 245	
 aag ttg gtt gcc gta agg gat cct cat ggg ttt agg cca ttg gtt atg	819
Lys Leu Val Ala Val Arg Asp Pro His Gly Phe Arg Pro Leu Val Met	
250 255 260 265	
 ggg agg aga agt aat ggt gct gtt gtt ttc gcg tct gag acg tgt gct	867
Gly Arg Arg Ser Asn Gly Ala Val Val Phe Ala Ser Glu Thr Cys Ala	
270 275 280	
 ttg gat ttg att gag gct act tat gag agg gag gtg aat cct ggt gag	915
Leu Asp Leu Ile Glu Ala Thr Tyr Glu Arg Glu Val Asn Pro Gly Glu	
285 290 295	
 gtt gtt gtt gtg gat aaa gat ggg gtt cag tct att tgt ttg atg cct	963
Val Val Val Val Asp Lys Asp Gly Val Gln Ser Ile Cys Leu Met Pro	
300 305 310	
 cat cct gag cgt aaa tct tgt atc ttt gag cat att tac ttt gct ctg	1011
His Pro Glu Arg Lys Ser Cys Ile Phe Glu His Ile Tyr Phe Ala Leu	
315 320 325	

[illegible]

## 10

tcg ggc aat tac cca gtc gag ccg acg ggt aag gtt aaa agg ata ggg 1635  
 Ser Gly Asn Tyr Pro Val Glu Pro Thr Gly Lys Val Lys Arg Ile Gly  
                   525                                  530                                  535

gat ttc atg gat gat gga tta agt gga gat atg gat tcc att gat ggt 1683  
 Asp Phe Met Asp Asp Gly Leu Ser Gly Asp Met Asp Ser Ile Asp Gly  
                   540                                  545                                  550

gga tgg cta cca gga agt agt agg gtt caa aag act atc ttg aat gaa 1731  
 Gly Trp Leu Pro Gly Ser Ser Arg Val Gln Lys Thr Ile Leu Asn Glu  
                   555                                  560                                  565

gtt aga acc agc taaactttct tttccatggt tgcttttagtt tttgcttttg 1783  
 Val Arg Thr Ser  
 570

atttctaattg cttgaccata gaaattataa gtttcaatga agtctctttt tctatttgga 1843

atgccacatg attctactga tctatg 1869

<210> 4

<211> 573

<212> PRT

<213> Nicotiana tabacum

<400> 4

Met Ala Ala Thr Val Ser Thr Ala Ser Ala Ala Ala Thr Asn Lys Tyr  
           1                                  5                                  10                                  15

Pro Leu Ser Gln Pro Leu Asp Lys Pro Phe Cys Ser Leu Ser Gln Lys  
                   20                                  25                                  30

Leu Leu Ser Leu Ser Pro Lys Thr His Pro Lys Pro Tyr Arg Thr Leu  
                   35                                  40                                  45

Ile Thr Ala Ser Ser Lys Asn Pro Leu Asn Asp Val Ile Ser Phe Lys  
           50                                  55                                  60

Lys Ser Ala Asp Asn Thr Leu Asp Ser Tyr Phe Asp Asp Asp Asp Lys  
           65                                  70                                  75                                  80

Pro Arg Glu Glu Cys Gly Val Val Gly Ile Tyr Gly Asp Ser Glu Ala  
                   85                                  90                                  95

Ser Arg Leu Cys Tyr Leu Ala Leu His Ala Leu Gln His Arg Gly Gln  
           100                                  105                                  110

## 11

Glu Gly Ala Gly Ile Val Ala Val Asn Asp Asp Val Leu Lys Ser Ile  
 115 120 125

Thr Gly Val Gly Leu Val Ser Asp Val Phe Asn Glu Ser Lys Leu Asp  
 130 135 140

Gln Leu Pro Gly Asp Met Ala Ile Gly His Val Arg Tyr Ser Thr Ala  
 145 150 155 160

Gly Ser Ser Met Leu Lys Asn Val Gln Pro Phe Val Ala Ser Tyr Lys  
 165 170 175

Phe Gly Ser Val Gly Val Ala His Asn Gly Asn Leu Val Asn Tyr Lys  
 180 185 190

Leu Leu Arg Ser Glu Leu Glu Glu Asn Gly Ser Ile Phe Asn Thr Ser  
 195 200 205

Ser Asp Thr Glu Val Val Leu His Leu Ile Ala Ile Ser Lys Ala Arg  
 210 215 220

Pro Phe Leu Leu Arg Ile Val Glu Ala Cys Glu Lys Ile Glu Gly Ala  
 225 230 235 240

Tyr Ser Met Val Phe Val Thr Glu Asp Lys Leu Val Ala Val Arg Asp  
 245 250 255

Pro His Gly Phe Arg Pro Leu Val Met Gly Arg Arg Ser Asn Gly Ala  
 260 265 270

Val Val Phe Ala Ser Glu Thr Cys Ala Leu Asp Leu Ile Glu Ala Thr  
 275 280 285

Tyr Glu Arg Glu Val Asn Pro Gly Glu Val Val Val Val Asp Lys Asp  
 290 295 300

Gly Val Gln Ser Ile Cys Leu Met Pro His Pro Glu Arg Lys Ser Cys  
 305 310 315 320

Ile Phe Glu His Ile Tyr Phe Ala Leu Pro Asn Ser Val Val Phe Gly  
 325 330 335

Arg Ser Val Tyr Glu Ser Arg Arg Ala Phe Gly Glu Ile Leu Ala Thr  
 340 345 350

Glu Ala Pro Val Glu Cys Asp Val Val Ile Ala Val Pro Asp Ser Gly  
 355 360 365

12

Val Val Ala Ala Leu Gly Tyr Ala Ala Lys Ala Gly Val Pro Phe Gln  
370 375 380

Gln Gly Leu Ile Arg Ser His Tyr Val Gly Arg Thr Phe Ile Glu Pro  
385 390 395 400

Ser Gln Lys Ile Arg Asp Phe Gly Val Lys Leu Lys Leu Ser Pro Val  
405 410 415

Arg Ala Val Leu Glu Gly Lys Arg Val Val Val Val Asp Asp Ser Ile  
420 425 430

Val Arg Gly Thr Thr Ser Ser Lys Ile Val Arg Leu Leu Lys Glu Ala  
435 440 445

Gly Ala Lys Glu Val His Met Arg Ile Ala Ser Pro Pro Ile Ile Ala  
450 455 460

Ser Cys Tyr Tyr Gly Val Asp Thr Pro Ser Ser Asp Glu Leu Ile Ser  
465 470 475 480

Asn Arg Met Ser Val Glu Glu Ile Lys Glu Phe Ile Gly Ser Asp Ser  
485 490 495

Leu Ala Phe Leu Pro Met Asp Ser Leu Asn Lys Leu Leu Gly Asn Asp  
500 505 510

Ser Lys Ser Phe Cys Tyr Ala Cys Phe Ser Gly Asn Tyr Pro Val Glu  
515 520 525

Pro Thr Gly Lys Val Lys Arg Ile Gly Asp Phe Met Asp Asp Gly Leu  
530 535 540

Ser Gly Asp Met Asp Ser Ile Asp Gly Gly Trp Leu Pro Gly Ser Ser  
545 550 555 560

Arg Val Gln Lys Thr Ile Leu Asn Glu Val Arg Thr Ser  
565 570